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20	30	40	50	
34567890	1234567890	1234567890	1234567890	
SAIYATSL	ASIIAFVAAG	CGQTESGSTS	DSKPQAETLK	50
TDPDNPRW	ISAOKDIISY	VDETEAATST	ITKNODAONN	100
KGFIIAPE	NGSGVGTAVN	TIADKGIPIV	AYDRLITGSD	150
VGELQGLS	LAAGLLGKED	GAFDSIDOMN	EYLKSHMPOF	200
DNNSQYFY	NGAMKVLKEL	MKNSONKIID	LSPEGENAVY	250
IQSFLTIN	KDPAGGNKIK	AVGSKPASIF	KGFLAPNDGM	300
FDTOKIFV	TRODYNDKAK	TEIKDGDONM	TIYKPDKVLG	350
KKNKASRS	EVENELKAKI	PNISEKYDNO	TYKVOGKNIN	400
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(57) Abstract

The present invention relates to a putative protective antigen against a Mycoplasma, prepared by a method including a sample of a Mycoplasma; an antibody probe including at least one antibody against a Mycoplasma produced by a method including: providing a biological sample taken a short time after an immune animal has been challenged with a Mycoplasma or Mycoplasma extract taken from the infection site or an area of a lesion or an area close to the infection site or lesion; isolating cells from the biological sample; culturing cells in vitro in a suitable culture medium; and harvesting antibodies produced from said cells; probing the Mycoplasma sample with the antibody probe to detect at least one antigen; and isolating the antigen detected, also including diagnostic antigens, the preparation thereof, and their use in the formation of vaccine compositions, particularly vaccine compositions against Mycoplasma hyopneumoniae infections.

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ANTIGEN COMPOSITION AGAINST MYCOPLASMA

The present invention relates to protective and diagnostic antigens, the preparation thereof, and their use in the formation of vaccine compositions, particularly vaccine compositions against Mycoplasma hyopneumoniae infections.

Mycoplasma hyopneumoniae is a ubiquitous swine respiratory pathogen causing mycoplasmal pneumoniae in swine (swine enzootic pneumonia). Swine enzootic pneumonia is probably the most widespread and economically significant disease in swine producing countries of the world. The economic effects of swine enzootic pneumonia (SEP) are complex, and the cost of the disease is severe. In Australia, the disease was estimated in 1988 to cost approximately \$20,000,000 per annum. Increased mortality, decreased growth weight, depressed feed conversion, susceptibility to secondary bacterial infections, increased management costs, and increased use of antibiotics, are the main reasons for the economic impact of SEP.

Whilst several experimental vaccines have been produced, these have resulted in less than optimal results, and utilising various classes of antibiotics such as tetracycline, lincamycin and tiamulin is still the most widespread control treatment. Such antibiotics are, however, of limited therapeutic value, because they do not prevent the establishment of an infection, and lung lesions may develop after treatment ends.

European Patent Application 359,919 to ML Technology Ventures L.P. describes a series of antigens, 36 kD, 41 kD, 74.5 kD and 96 kD in size, and proposes the use of such antigens in vaccines. Results presented suggest that some protection in pigs against challenge was achieved.

However, there remains a need in the art for an effective vaccine against M. hyopneumoniae which would confer protection against colonisation and clinical disease following M. hyopneumoniae challenge and also significantly reduce the morbidity and mortality from secondary infections.

Accordingly, it is an object of the present invention to overcome, or at least alleviate, one or more of the difficulties and deficiencies in the prior art.

Accordingly, in a first aspect of the present invention there is provided a putative protective antigen against a <u>Mycoplasma</u>, preferably <u>Mycoplasma</u>

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hyopneumoniae prepared by a method including providing

a sample of a Mycoplasma;

an antibody probe including at least one antibody against a Mycoplasma produced by a method including;

providing a biological sample taken a short time after an immune animal has been challenged with a <u>Mycoplasma</u> or <u>Mycoplasma</u> extract taken from the infection site or an area of a lesion or an area close to the infection site or lesion:

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isolating cells from the biological sample; culturing cells <u>in vitro</u> in a suitable culture medium; and harvesting antibodies produced from said cells;

probing the <u>Mycoplasma</u> sample with the antibody probe to detect at least one antigen; and

isolating the antigen detected.

The protective antigens may also function as diagnostic antigens as discussed below.

Accordingly, in a preferred aspect of the present invention there is provided a putative protective antigen against <u>Mycoplasma hyopneumoniae</u>, or related infections, selected from the group of antigens having approximate molecular weights of 110-114, 90-94, 72-75, 60-64, 52-54 and 46-48 kilodaltons (kD), as hereinafter described, mutants, derivatives and fragments thereof. The putative protective antigen may be a surface protein. The putative protective antigen may be a surface lipoprotein or membrane protein.

Preferably the protective antigens are selected from the group of antigens having approximate molecular weights of 110-114, 90-94, 74, 62, 52 and 48 kD.

Preferably, the 72-75 kD antigen includes the following N-terminal amino acid sequence:

AGXLQKNSLLEEVWYLAL

and, optionally, one or more of the following internal amino acid sequences:

AKNFDFAPSIQGYKKIAHEL NLKPEQILQLLG

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LLKAEXNKXIEEINTXLDN

Preferably, the 60-64 kD antigen includes one of the following N-terminal amino acid sequences:

MKLAKLLKGFX(N/L)(M/V)IK

5 ADP(F/I)(R/E)Y(V/A)PQG(Q/A)X(M/N)VG

Preferably, the 52-54 kD antigen includes the following N-terminal amino acid sequence:

AGXWAKETTKEEKS

and, optionally, one or more of the following internal amino acid sequences:

10 AWVTADGTVN

AIVTADGTVNDNKPNQWVRKY.

Preferably, the 46-48 kD antigen includes the following N-terminal amino acid sequence:

AGXGQTESGSTSDSKPQAETLKHKV

and, optionally, one or more of the following internal amino acid sequences:

TIYKPDKVLGKVAVEVLRVLIAKKNKASR

AEQAITKLKLEGFDTQ

KNSQNKIIDLSPEG

The 46-48 kD antigen may be encoded by a nucleic acid fragment:

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10	20	30	40	50	
1234567890	1234567890	1234567890	1234567890	1234567890	
ATGAAAAAA	TGCCACTATA	CCAGAGGAAA	GAGCAGTATA	TAAAATAATT	50
AAAATTACAT	TTTCTTCATT	TGCGCCAGAA	TTTTTAAGAA	TTAGTACATT	100
AAAAAGTAGA	ACAAAAGTTA	TTAATGTAAA	CATTAGCGCA	ATCCTTAAGA	150
AAAAATTAAA	AGTTTTATCT	ATTTTTTA	ATCGAAATCC	AACCAGGCAT	200
AAATCTTTGT	CAGTATTTAT	CAAGTCGGTA	TTTTTCATT	ATTTCTACTA	250
TTATTATA	TGAATTTGCA	TTTTCCATAA	TCTAAAATTT	TACATTTTTT	300
TATAACAATT	TTAAAAATT	ACTCTTTAAT	TTATAGTATT	TTTTATTTT	350
TTAGTCTAAA	TTAAAAATT	ATCTTGAATT	TTATTTGAAT	TTTATAATT	400
TAGTACTAAA	AAATACAAAT	ATTTTTCCT	ATTCTAAGAA	AAATTCATTT	450
TTTAAAAAAA	ATTGATTTTT	ATAGTATAAT	TTGTTTGTAT	AATTGAATTA	500
ACTTGATTTG	AAAGGGAACA	AAATGAAAAA	AATGCTTAGA	AAAAAATTCT	550
TGTATTCATC	AGCTATTTAT	GCAACTTCGC	TTGCATCAAT	TATTGCATTT	600
GTTGCAGCAG	GTTGTGGACA	GACAGAATCA	GGTTCAACTT	CTGATTCTAA	650
ACCACAAGCC	GAGACGCTAA	AACATAAAGT	AAGTAATGAT	TCTATTCGAA	700
TAGCACTAAC	CGATCCGGAT	AATCCTCGAT	GAATTAGTGC	CCAAAAAGAT	750
ATTATTTCTT	ATGTTGATGA	AACAGAGGCA	GCAACTTCAA	CAATTACAAA	800
AAACCAGGAT	GCACAAAATA	ACTGACTCAC	TCAGCAAGCT	AATTTAAGCC	850
CAGCGCCAAA	AGGATTTATT	ATTGCCCCTG	AAAATGGAAG	TGGAGTTGGA	900

	ACTGCTGTTA	ATACAATTGC	TGATAAAGGA	ATTCCGATTG	TTGCCTATGA	950
	TCGACTAATT	ACTGGATCTG	ATAAATATGA	TTGGTATGTT	TCTTTTGATA	1000
	ATGAAAAAGT	TGGTGAATTA	CAAGGTCTTT	CACTTGCTGC	GGGTCTATTA	1050
	GGAAAAGAAG	ATGGTGCTTT	TGATTCAATT	GATCAAATGA	ATGAATATCT	1100
5	AAAATCACAT	ATGCCCCAAG	AGACAATTTC	TTTTTATACA	ATCGCGGGTT	1150
	CCCAAGATGA	TAATAATTCC	CAATATTTTT	ATAATGGTGC	AATGAAAGTA	1200
	CTTAAAGAAT	TAATGAAAAA	TTCGCAAAAT	AAAATAATTG	ATTTATCTCC	1250
	TGAAGGCGAA	AATGCTGTTT	ATGTCCCAGG	ATGAAATTAT	GGAACTGCCG	1300
	GTCAAAGAAT	CCAATCTTTT	CTAACAATTA	ACAAAGATCC	AGCAGGTGGT	1350
10	AATAAAATCA	AAGCTGTTGG	TTCAAAACCA	GCTTCTATTT	TCAAAGGATT	1400
	TCTTGCCCCA	AATGATGGAA	TGGCCGAACA	AGCAATCACC	AAATTAAAAC	1450
	TTGAAGGGTT	TGATACCCAA	AAAATCTTTG	TAACTCGTCA	AGATTATAAT	1500
	GATAAAGCCA	AAACTTTTAT	CAAAGACGGC	GATCAAAATA	TGACAATTTA	1550
	TAAACCTGAT	AAAGTTTTAG	GAAAAGTTGC	TGTTGAAGTT	CTTCGGGTTT	1600
15	TAATTGCAAA	GAAAAATAAA	GCATCTAGAT	CAGAAGTCGA	AAACGAACTA	1650
	AAAGCAAAAC	TACCAAATAT	TTCATTTAAA	TATGATAATC	AAACATATAA	1700
	AGTACAAGGT	AAAAATATTA	ATACAATTTT	AGTAAGTCCA	GTAATTGTTA	1750
	CAAAAGCTAA	TGTTGATAAT	CCTGATGCCT	AA		1782

Accordingly, in a further aspect the present invention provides an isolated nucleic acid fragment encoding a putative protective antigen against Mycoplasma hyopneumoniae or related infections, said nucleic acid fragment:

	10	20	30	40	50	
25	1234567890	1234567890	1234567890	1234567890	1234567890	
		······································				
	ATGAAAAAA	TGCCACTATA	CCAGAGGAAA	GAGCAGTATA	TAAAATAATT	50
	AAAATTACAT	TTTCTTCATT	TGCGCCAGAA	TTTTTAAGAA	TTAGTACATT	100
	AAAAAGTAGA	ACAAAAGTTA	TTAATGTAAA	CATTAGCGCA	ATCCTTAAGA	150
30	AAAAATTAAA	AGTTTTATCT	ATTTTTTTA	ATCGAAATCC	AACCAGGCAT	200
	AAATCTTTGT	CAGTATTTAT	CAAGTCGGTA	TTTTTCATT	ATTTCTACTA	250
	TTATTATA	TGAATTTGCA	TTTTCCATAA	TCTAAAATTT	TACATTTTTT	300
	TATAACAATT	TTAAAAATT	ACTCTTTAAT	TTATAGTATT	TTTTATTTT	350
	TTAGTCTAAA	TTATAAAATT	ATCTTGAATT	TTATTTGAAT	TTTATAATT	400
35	TAGTACTAAA	AAATACAAAT	ATTTTTCCT	ATTCTAAGAA	AAATTCATTT	450
	TTTAAAAAAA	ATTGATTTTT	ATAGTATAAT	TTGTTTGTAT	AATTGAATTA	500
	ACTTGATTTG	AAAGGGAACA	AAATGAAAAA	AATGCTTAGA	AAAAAATTCT	550
	TGTATTCATC	AGCTATTTAT	GCAACTTCGC	TTGCATCAAT	TATTGCATTT	600
	GTTGCAGCAG	GTTGTGGACA	GACAGAATCA	GGTTCAACTT	CTGATTCTAA	650
40	ACCACAAGCC	GAGACGCTAA	AACATAAAGT	AAGTAATGAT	TCTATTCGAA	700
	TAGCACTAAC	CGATCCGGAT	AATCCTCGAT	GAATTAGTGC	CCAAAAAGAT	750
	ATTATTTCTT	ATGTTGATGA	AACAGAGGCA	GCAACTTCAA	CAATTACAAA	800
	AAACCAGGAT	GCACAAAATA	ACTGACTCAC	TCAGCAAGCT	AATTTAAGCC	850
	CAGCGCCAAA	AGGATTTATT	ATTGCCCCTG	AAAATGGAAG	TGGAGTTGGA	900
45	ACTGCTGTTA	ATACAATTGC	TGATAAAGGA	ATTCCGATTG	TTGCCTATGA	950
	TCGACTAATT	ACTGGATCTG	ATAAATATGA	TTGGTATGTT	TCTTTTGATA	1000
	ATGAAAAAGT	TGGTGAATTA	CAAGGTCTTT	CACTTGCTGC	GGGTCTATTA	1050
	GGAAAAGAAG	ATGGTGCTTT	TGATTCAATT	GATCAAATGA	ATGAATATCT	1100
	AAAATCACAT	ATGCCCCAAG	AGACAATTTC	TTTTTATACA	ATCGCGGGTT	1150
50	CCCAAGATGA	TAATAATTCC	CAATATTTTT	ATAATGGTGC	AATGAAAGTA	1200
	CTTAAAGAAT	TAATGAAAAA	TTCGCAAAAT	AAAATAATTG	ATTTATCTCC	1250
	TGAAGGCGAA	AATGCTGTTT	ATGTCCCAGG	ATGAAATTAT	GGAACTGCCG	1300
	GTCAAAGAAT	CCAATCTTTT	CTAACAATTA	ACAAAGATCC	AGCAGGTGGT	1350

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AATAAAATCA	AAGCTGTTGG	TTCAAAACCA	GCTTCTATTT	TCAAAGGATT	1400
TCTTGCCCCA	AATGATGGAA	TGGCCGAACA	AGCAATCACC	AAATTAAAAC	1450
TTGAAGGGTT	TGATACCCAA	AAAATCTTTG	TAACTCGTCA	AGATTATAAT	1500
GATAAAGCCA	AAACTTTTAT	CAAAGACGGC	GATCAAAATA	TGACAATTTA	1550
TAAACCTGAT	AAAGTTTTAG	GAAAAGTTGC	TGTTGAAGTT	CTTCGGGTTT	1600
TAATTGCAAA	GAAAAATAAA	GCATCTAGAT	CAGAAGTCGA	AAACGAACTA	1650
AAAGCAAAAC	TACCAAATAT	TTCATTTAAA	TATGATAATC	AAACATATAA	1700
AGTACAAGGT	ATATAAAAA	ATACAATTTT	AGTAAGTCCA	GTAATTGTTA	1750
CAAAAGCTAA	TGTTGATAAT	CCTGATGCCT	AA		1782
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As cross protection between various <u>Mycoplasma</u> such as <u>M. hyorhinis</u> and <u>M. synoviae</u> has been documented, similar antigens may also be detected in other <u>Mycoplasma</u> species (Figure 1).

In a still further aspect the present invention provides a method for preventing Mycoplasma infection in animals. Preferably the Mycoplasma disease is a Mycoplasma hyopneumoniae disease such as swine enzootic pneumonia (SEP). This method includes administering to an animal an effective amount of at least one protective antigen against Mycoplasma as described above.

The present invention further provides a vaccine composition including a prophylactically effective amount of at least one putative protective antigen against a Mycoplasma as herein described. Preferably the veterinary composition includes two or more putative protective antigens as herein described.

Accordingly in a preferred aspect the present invention provides a vaccine composition including two or more putative protective antigens selected from antigens having approximate molecular weights of 110-114, 90-94, 72-75, 60-64, 52-54 and 46-48 kilodaltons.

The vaccine composition may include any combination of two or more putative protective antigens selected from antigens having approximate molecular weights of 110-114, 90-94, 72-75, 60-64, 52-54 and 46-48 kD. The two or more antigens may be selected from antigens falling within one of the specified approximate molecular weights and/or antigens from different specified approximate molecular weights. The composition may contain 3, 4, 5 or 6 antigens selected from protective antigens having molecular weights of approximately 110-114, 90-94, 72-75, 60-64, 52-54 and 46-48 kD.

The vaccine compositions according to the present invention may be

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administered orally or may be administered parenterally (for example by intramuscular, subcutaneous, intradermal or intravenous injection). The amount required will vary with the antigenicity of the active ingredient and need only be an amount sufficient to induce an immune response typical of existing vaccines.

Reactive experimentation will easily establish the required amount. Typical initial doses of vaccine or veterinary compositions may be approximately 0.001-1 mg active ingredient/kg body weight. The dose rate may increase or multiple doses may be used as needed to provide the desired level of protection.

The vaccine composition according to the present invention may further include a veterinary acceptable carrier, diluent or excipient therefor. Preferably the active ingredient may be suspended or dissolved in a carrier. The carrier may be any solid or solvent that is nontoxic to the animal and compatible with the active ingredient. Suitable carriers include liquid carriers, such as normal saline and other nontoxic salts at or near phsyiological concentrations, and solid carriers, such as talc or sucrose.

Preferably the vaccine contains an adjuvant, such as Freund's adjuvant, complete or incomplete, or immunomodulators such as cytokines may be added to enhance the antigenicity of the antigen if desired.

More preferably the adjuvant is of the mineral-oil type as these have been found to be consistently superior at inducing antibody titres and Delayed Type Hypersensitivity responses. A particularly preferred adjuvant is that marketed under the trade designation Montanide ISA-50 and available from Seppic, Paris, France.

When used for administering via the bronchial tubes, the vaccine is suitably present in the form of an aerosol.

In a still further aspect of the present invention there is provided a diagnostic kit including a diagnostic antigen against a <u>Mycoplasma</u>, preferably <u>Mycoplasma</u> <u>hypneumoniae</u>, identified and purified as described above.

The putative protective antigens according to the present invention may be isolated and identified utilising the general methods described in Australian patent application 49035/90, the entire disclosure of which is incorporated herein by reference.

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Accordingly, in a further aspect, the present invention provides a method for producing at least one antibody against a <u>Mycoplasma</u>. This method includes

providing a biological sample taken a short time after an immune animal has been challenged with a <u>Mycoplasma</u> or <u>Mycoplasma</u> extract taken from the infection site or an area of a lesion or an area close to the infection site or lesion;

isolating cells from the biological sample;

culturing cells <u>in vitro</u> in a suitable culture medium; and harvesting antibodies produced from said cells.

The Mycoplasma may be Mycoplasma hyopneumoniae.

The animal may be a mammal including humans. The mammal may be a domestic animal such as a pig, sheep or cattle.

The biological animal sample may be of any suitable type. The biological sample may be taken from animal tissue, organs, lymph or lymph nodes. The biological sample may be taken from the infection site, the lungs of the animal, or an area of a lesion which may be formed or an area close to the infected site or a lesion such as in the lymph nodes draining from the lungs.

However, serum/plasma samples are not used as the biological samples according to this aspect of the present invention. It has been found that the majority of antibodies found in a serum/plasma sample are irrelevant to protection or specific diagnosis or a Mycoplasma or are unrelated to the Mycoplasma. In addition, other serum/ plasma components may interfere with the specific reactions between pathogen components and antibodies to them.

In contrast, the probes described in the present invention are highly enriched in <u>Mycoplasma</u>-specific antibodies of particular importance to protective immunity.

It is preferred that the biological samples are taken from the animals at a predetermined time in the development of the disease. In general, for a Mycoplasma infection, it has been found that the biological samples should be taken approximately 2 to 7 days after challenge with or after administration of products obtained from a pathogen or with the pathogen itself.

The cells isolated from the biological sample may include B cells.

Thus, preferably the cells are taken a short time after in vivo stimulation,

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preferably within approximately 2 to 5 days thereafter, resulting in the <u>in vivo</u> induction of antibody forming cells which will secrete specific antibodies into the culture medium after <u>in vitro</u> incubation.

In vitro secretion of antibodies in the culture medium by recently activated B cells may be enhanced by the addition of helper factors to the cultures. The helper factors may be cytokines used alone or in combination, including Interleukin 1, 2, 3, 4, 5, 6, 7 and 8, colony stimulating factors, interferons and any other factors that may be shown to have an enhancing effect on specific B cell secretion.

The method of producing an antibody may include a further step of activating the cells isolated to proliferate and secrete and/or release antibodies.

The cell activation step may include adding a cell activating agent to the culture medium. The cell activating agent may be selected from mitogens and helper factors produced by leukocytes, or their synthetic equivalents or combinations thereof.

The mitogens may be selected from the group including products derived from pokeweed (Phytolacca americana) also known as pokeweed mitogen (PWM), polyvinylpyrrolidone (PVP), polyadenylic-polyuridylic acid (poly(A-U)), purified protein derivate (PPD), polyinosinic-polycytidilic acid (poly(I-C)), lipopolysaccharide (LPS), staphylococcal organisms or products thereof, Bactostreptolysin O reagent (SLO), Staphylococcal phage lysate (SPL), Epstein-Barr virus (EBV). Nocardia water-soluble mitogen (NWEM), phytohemagglutinin (PHA), Concanavalin A (Con A), and dextran-sulphate and mixtures thereof. The cell proliferation agent may be any agent that indirectly or directly results in B cell proliferation and/or antibody secretion such as solid-phase anti-immunoglobulin. The helper factors may be selected from the group including cytokines including interleukin 1, 2, 3, 4, 5, 6, 7 and 8, colony stimulating factors, interferons and any other helper factors that may be shown when added alone, or in combination with other factors and agents, to have an enhancing effect on specific B cell proliferation and/or antibody secretion. This in no way is meant to be an exhaustive list of mitogens and cell actuating agents including helper factors.

The in vitro culturing of the cells may be conducted with or without prior

steps to separate sub-populations of cells. The harvesting of antibodies may be conducted by harvesting of the supernatant from the culture medium. This supernatant contains antibodies secreted by these cells during the <u>in vitro</u> culture or artificially released from the B cells, for example by lysis of the B cells. It has been found that the antibody-containing supernatants may be used directly to detect antigens of the <u>Mycoplasma</u>.

In a preferred aspect of the present invention, there is provided a method for identifying an antigen associated with a Mycoplasma, preferably Mycoplasma, preferably Mycoplasma, <a href="Mycoplas

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a sample of a Mycoplasma; and

an antibody probe including at least one antibody against a Mycoplasma;

probing the <u>Mycoplasma</u> sample with the antibody probe to detect at least one antigen; and

isolating the antigen detected.

The sample of <u>Mycoplasma</u> may be mixed with a standard buffer solution and placed on a standard support such as an SDS-polyacrylamide gel to separate the proteins contained thereon (Figure 2).

Alternatively, the proteins may be selected utilising the non-ionic detergent Triton X-114 (TX-114). Insoluble material may be removed by centrifugation. Proteins soluble in the TX-114 phase may then be precipitated out (Figure 2).

The separate proteins may then be transferred to nitrocellulose, nylon or other sheets.

The probing with a suitable antibody may further include subjecting the product produced thereby to a detection assay. The detection assay may include Western blot techniques. The detection assay may be an immunoprecipitation assay, a radioimmunoassay, an enzyme-linked immunoassay or immunofluorescent assay (Figures 3, 4 and 5).

The antibody produced as described above may be utilized simply in the form of the supernatant harvested from the culture medium. Alternatively, the antibodies may be separated and purified.

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In a further preferred aspect of the present invention the antibody contained in the culture medium may be used for the affinity purification, preferably immuno-affinity purification of antigen.

Accordingly, in a preferred aspect there is provided a method for purifying antigen. This method includes

providing

a crude antigen mixture; and

an antibody against a <u>Mycoplasma</u> immobilized on a suitable support;

subjecting the crude antigen mixture to affinity chromatography utilizing the immobilized antibody; and

isolating the purified antigen so formed.

The antibody is produced by the method described above.

Antibody can be obtained from the culture supernatant probe by conventional methods. For example, methods usually used to purify immunoglobulins from serum or plasma, e.g. precipitation with ammonium sulphate, fractionation with caprylic acid, ion exchange chromatography, or by binding and elution from immobilized protein G or protein A, may be utilized. Antibody so obtained can then be coupled to suitable supports, e.g., CNBractivated Sepharose 4B (Pharmacia), Affi-gel (Bio-RAD), or other affinity chromatography supports able to bind proteins.

Immobilized antibody can then be applied to the fractionation and purification of specific antigen from a complex Mycoplasma extract by affinity chromatography. After binding of antigen to immobilized antibody, unbound macromolecular species can be washed away from the solid support with, e.g. buffers containing 1.5 M NaCl. Subsequently the antigen can be eluted from the affinity column with, e.g. low or high pH buffer or buffers containing chaotropic ions, e.g. 0.5-3.0 M sodium thiocyanate.

The application of the antibody probe to affinity chromatography enables sufficient quantities of specific antigens to be rapidly isolated from a complex crude extraction mixture for biochemical characterization, amino-acid sequencing and vaccination of animal for limited protection studies. Application of affinity

chromatography for obtaining antigen(s) avoids the difficulties often encountered when applying conventional biochemical techniques to the purification of an antigen about which little or no data is known. It also obviates the need to raise polyclonal or monoclonal antibodies for the purpose of "analytical" affinity chromatography. Large scale preparation may, however, require the preparation of polyclonal or monoclonal antibodies.

Having identified the antigen(s) molecular biology, chemical techniques, e.g. cloning techniques, may be used to produce unlimited amounts of this antigen or, alternatively, synthetic peptides corresponding to different fragments of the identified antigens may be used as a means to produce a vaccine.

Accordingly in a preferred aspect of the present invention there is provided a method for preparing a synthetic antigenic polypeptide against <u>Mycoplasma</u>, preferably <u>Mycoplasma</u> <u>hypneumoniae</u>, which method includes

providing

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a cDNA library or genomic library derived from a sample of Mycoplasma; and

an antibody probe as described above;

generating synthetic polypeptides from the cDNA library or genomic library; probing the synthetic polypeptides with the antibody probe; and isolating the synthetic antigenic polypeptide detected thereby.

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Either cDNA or genomic libraries may be used. The cDNA or genomic libraries may be assembled into suitable expression vectors that will enable transcription and the subsequent expression of the clone cDNA, either in prokaryotic hosts (e.g. bacteria) or eukaryotic hosts (e.g. mammalian cells). The probes may preferably be selected from

- synthetic oligonucleotide probes based on the amino acid sequence of the antigen identified and purified as described above;
- (ii) antibodies obtained from the culture medium produced as described above;
- 30 (iii) monoclonal or polyclonal antibodies produced against the antigens identified and purified as described above;
 - (iv) recombinant or synthetic monoclonal antibodies or polypeptides with

specificity for the antigen, e.g. as described by Ward et al., <u>Nature</u>, <u>241</u>, pages 544-546 (1989).

The synthetic antigenic polypeptide produced in accordance with the invention may be a fusion protein containing the synthetic antigenic peptide and another protein.

In a further aspect of the present invention there is provided a DNA fragment encoding a putative protective antigen against Mycoplasma or related infections, said DNA fragments having a nucleic acid sequence according to Figure 6a and 6b or an homologous sequence and functionally active fragments thereof.

In a further preferred aspect of the present invention there is provided a clone including a DNA fragment encoding a putative protective antigen against Mycoplasma or related infections, said DNA fragments having a nucleic acid sequence according to Figure 6a and 6b or an homologous sequence and functionally active fragments thereof.

Preferably the clone is pC1-2.

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The present invention will now be more fully described with reference to the accompanying Examples and drawings. It should be understood, however, that the description following is illustrative only and should not be taken in any way as a restriction on the generality of the invention described above.

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IN THE FIGURES:

<u>FIGURE 1</u>: SDS-Polyacrylamide gel (12.5%) profiles of SDS extracts of species of mycoplasma- Coomassie R250 stained.

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- Lane 1 Pre-stained Molecular Weight Standards.
- Lane 2 M. gallisepticum.

Lane 3 M. synoviae.

Lane 4 M. hyopneumoniae.

Lane 5 M. hyorhinis.

Lane 6 M. flocculare.

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FIGURE 2: SDS-Polyacrylamide gel (12.5%) profiles of extracts of strains of M. hyopneumoniae - Coomassie R250 stained gel

Lane 1 Pre-stained Molecular Weight Standards.

10 Lane 2 Triton X-114 extract of M. hyopneumoniae - strain Beaufort.

Lane 3 As for Lane 2.

Lane 4 SDS extract of *M. hyopneumoniae* strain Beaufort.

Lane 5 SDS extract of *M. hyopneumoniae* strain 10110.

- 15 <u>FIGURE 3</u>: Western blots of Triton X-114 extracted antigens from *M. hyopneumoniae* strain Beaufort, probed with serum and supernatant antibody probes.
 - Lane 1 No antibody control.
- 20 Lane 2 Dookie pig serum control 1/200.
 - Lane 3 Pig 105 supernatant.
 - Lane 4 Pig 1 supernatant.
 - Lane 5 Dookie pig supernatant.
- 25 <u>FIGURE 4</u>: Western blots of SDS extracted antigens from *M. hyopneumoniae* strain Beaufort probed with paired serum and supernatantantibody probes. Fractionation of antigens on SDS Polyacrylamide gel (12.5%).
 - Lane 1 a) Pig 453 supernatant.
- 30 b) Pig 453 serum 1/100.
 - Lane 2 a) Pig 105 supernanant.
 - b) Pig 105 serum 1/100.

- Lane 3 a) Pig 1 superanatant. b) Pig 1 serum 1/100.
- Lane 4 a) Pig 15 supernanant. b) Pig 15 serum 1/100.
- 5 Lane 5 a) Dookie supernatant.
 - b) Dookie serum 1/100.
 - Lane 6 No antibody control.

FIGURE 5: Western blots of SDS extracted antigens from *M. hyopneumoniae*strain Beaufort probed with paired serum and supernatantantibody probes.
Fractionation of antigens on SDS Polyacrylamide gel (10.0 %).

- Lane 1 a) Pig 453 supernatant.
 - b) Pig 453 serum 1/100.
- 15 Lane 2 a) Pig 105 supernatant.
 - b) Pig 105 serum 1/100.
 - Lane 3 a) Pig 1 supernatant.
 - b) Pig 1 serum 1/100.
 - Lane 4 a) Pig 15 supernatant.
- 20 b) Pig 15 serum 1/100.
 - Lane 5 a) Dookie supernatant.
 - b) Dookie serum 1/100.
 - Lane 6 No antibody control.
- 25 FIGURE 6: The entire 48 k gene sequence.
 - FIGURE 7: tHE 48kDa protein sequence of the 48k gene sequence.

EXAMPLE 1

Mycoplasma hyopneumoniae media

Friss Media

Hovind-Hougen, K., Friss, N.F., Research in Veterinary Science, 1991, 51, pp 155-163, "Morphological & Ultrastructural Studies of M flocculare and M hyppneumoniae in vitro".

250 ml Hanks BSS

10 140 ml Water

1.5 gm Brain Heart infusion

1.6 gm PPLO Broth w/o CV

Autoclave at 120°C for 20 minutes

18 ml Yeast Extract (100g YSC-2 Sigma in 750 ml)

15 3.7 ml 0.2% DNA in 0.1% Na₂CL₃

5.14 ml 1% -NAD

0.6 ml 1% Phenol red

Adjust to pH 7.3 to 7.4

Filter through 0.45 um, 0.2 um membrane, store at 4°C.
Add sterile Horse or Pig serum to 20%
and Antibiotics prior to use

Etheridge Media

Etheridge, J.R., Cottew, G.S., Lloyd, L.C., Australian Veterinary Journal, 1979, August <u>55</u>, pp 356-359, "Isolation of <u>Mycoplasma hyopneumoniae</u> from lesions in experimentally infected pigs".

	<u>Materials</u>	For 600 mls
	Hanks BSS	18.9 ml
	Hartleys Digest broth	1.28 gm
5	Heart Infusion broth	1.65 gm
	Lactalbumin hydrolysate	2.21 gm
	Glucose	4.41 gm
	Yeast Extract autolysate	8.82 ml
	Pig Serum (filtered)	163 ml
10	1% NAD	6.17 ml
	1% Phenol red	1.32 ml
	0.2% DNA in 0.1% Na ₂ CO ₃	4.41 ml

Make up to 600 ml with MQ water (about 350 - 400 ml)

Adjust pH to 7.4 and filter through: 3.0 um, 0.8 um, 0.45 um, 0.2 um.

Store at 4°C.

Development of Immune Sows

Cull sows and naive gilt (unmated sow designated Dookie).

20 Challenged on numerous occasions, with culture grown M. hyopneumoniae and lung homogenate. Given intranasally and intratracheally. Period of challenge - from September, 1991 to 21st January, 1992.

Tiamulin antibiotic given 31st January, 1992 to 4th February, 1992. Rested for approximately 8 weeks.

25 Infectious Challenge

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120 ml of frozen culture of <u>M. hyopneumoniae</u> strain Beaufort, spun down (12,000 xg, 20 min.) and resuspended in 50 ml complete medium and cultured overnight at 37°C. The overnight culture was centrifuged (12,000 xg, 20 min.) and the <u>Mycoplasma</u> cells resuspended in 10 ml serum free <u>Mycoplasma</u> culture medium. The 10 ml of concentrated mycoplasma was administered to anaesthetised immune sows via a catheter to ensure the inoculum was placed into the trachea.

Three of four days post-challenge, the sows were killed, and lymph nodes draining the lungs taken - these included the left and right tracheobronchial lymph nodes, and the lymph nodes located at the bifurcation of the trachea.

Antibody probes were prepared from pig lymph nodes and utilised to detect putative protection antigens as described in Australian Patent Application 49035/90 referred to above. Separate cell cultures were obtained from individual lymph nodes. Culture supernatants were harvested after 5 days of culture.

Antigen Preparation

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Mycoplasma hyopneumoniae strain Beaufort was cultured in Etheridge media until the pH had dropped to between 6.8 and 7.0. Cells of M. hyopneumoniae were harvested from culture by centrifugation at 12,000 xg for 20 min., washed 4 times with either sterile PBS or 0.25 M NaCl and then the pelleted cells extracted with one of the following.

(i) Sodium dodecyl sulphate (SDS)

The cell pellet was resuspended in 0.2% SDS and extracted for 2 hours at 37°C. Insoluble material was pelleted from the extract at 12,000 xg for 10 min. and the soluble extract run on SDS-polyacrylamide gel electrophoresis (SDS-PAGE).

(ii) Triton X-114

20 The method of Bordier (J. Bio. Chem. 1981, 256:1604-1606) was used to selectively extract membrane proteins using the non-ionic detergent Triton X-114.

The cell pellet was resuspended in cold PBS to 2 mg/ml protein and a cold pre-condensed solution of TX-114 added to give a final concentration of 1% (v/v) TX-114. Extraction was achieved by incubation overnight at 4°C with gentle mixing. Insoluble material was removed by centrifugation at 12,000 xg for 20 min. at 4°C. The Triton X-114 soluble membrane proteins were then obtained by achieving a phase separation at 37°C.

Proteins soluble in TX-114 phase were precipitated with 80% ethanol in the presence of carrier dextran (80,000 molecular weight) at -70°C overnight. The proteins were collected by centrifugation at 12,000 xg for 30 min. and dissolved to 500 ug/ml in 4 M urea.

Identification of Antigens

Six antigens were identified utilising the above- mentioned technique. The identified antigens were those that were consistently identified by the antibody probes from the immune cultures and the Dookie gilt. The results are summarised in Table 1.

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		TABLE 1
	Molecular Weight (kD)	Characteristics
	110-114	SDS Extracted
	90-94	SDS Extracted
10	72-75	Triton X-114 Extracted
	60-64**	SDS Extracted. Partitions to aqueous
		phase of Triton X-114 extract.
	52-54	Triton X-114 Extracted
	46-48	Triton X-114 Extracted

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^{**} Two antigens of approximate molecular weight 62 kD were identified.

	Molecular	Amino Acid Sequence
	Weight (kD)	
20		
	46-48	48 K N-Terminal: AGXGQTESGSTSDSKPQAETLKHKV
		48 K CNBR F 1: TIYKPDKVLGKVAVEVLRVLIAKKNKASR
		48 K CNBR F 2: AEQAITKLKLEGFDTQ
		48 K CNBR F 3: KNSQNKIIDLSPEG
25		
	52-54	52 K N-Terminal: AGXWAKETTKEEKS
		52 K CNBR F 1: AWVTADGTVN
		52 K CNBR F 2: AIVTADGTVNDNKPNQWVRKY
30	60-64	62 K N-Terminal: MKLAKLLKGFX (N/L)(M/V) IK
	60-64	62 K N-Terminal ADP(F/I)(R/E)Y(V/A)PQG(Q/A)X(M/N)VG

WO 96/28472

72-75 74 K N-Terminal: AGXLQKNSLLEEVWYLAL

74 K CNBR F 1: AKNFDFAPSIQGYKKIAHEL

74 K CNBR F 2: NLKPEQILQLLG

5 74 K CNBR F 3: LLKAEXNKXIEEINTXLDN

CNBR - Cyanogen Bromide fragment

X denotes an undetermined amino acid

(A/B) - residue may be A or B

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PCR of 48kDa Gene

Polymerase Chain Reaction (PCR) oligonucleotide primers were designed from the amino acid sequences obtained from the N-terminal and internal cyanogen bromide (CNBr) derived peptides. Inosine (I) was substituted at positions of high redundancy. The following primers were used in a standard PCR assay, run on a Bartelt Gene Machine Robotic thermal cycling instrument.

Oligo 48 K CNBr F 1: ACIAACGACGAGAAGCCICAGGC

TTAAA

20 Oligo 48 K CNBr F 2: TTIAGCTTIGTGATIGCCTGCTC

AT A T T

T

Oligo 48 K CNBr F 3: AGGTCGATGATCTTCCAICC

AA A A T T

TT

The resulting PCR products were visualised on a 1.5% agarose gel, excised, and purified using Prep-a-Gene (BioRad). They were cloned by standard techniques into a dideoxy tailed T-vector (Holton and Graham, Nucleic Acids Research 19: 1156, 1991) and the nucleic acid sequence determined. The PCR product, obtained from the reaction using primers F1 and F2 shown above, was of

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approximately 810 base pairs and was shown by sequencing to code for the previously determined amino acid sequence of the purified native 46-48kDa protein.

5 Genomic clone isolation at 48 k gene

The entire 48k gene and 48kDa protein (Figures 6 and 7) has been isolated and sequenced. The gene was obtained from an *M. hyopneumoniae* genomic library made by digesting genomic DNA with the restriction enzyme CLA I and ligating the fragments into the vector pBluescript (Stratagene). The ligated product was then electroporated into *Escherichia coli* strain SURE (Stratagene) and the cells plated on Luria Broth agar plates containing 100 µg/ml Ampicillin (LB-Amp). The library was screened by DNA hybridisation with a polymerase chain reaction (PCR) product specific for the 48 kDa protein. Positive clones were grown in LB-Amp, the cells harvested and the DNA isolated and partially sequenced for confirmation.

The positive clone pC1-2 was entirely sequenced and the protein sequence deduced. This was compared to the protein sequence obtained from the N terminus and Cyanogen Bromide fragments of the 48 kDa protein to show the that the gene encoded the desired protein.

Adjuvant Selection

Young piglets, 5-7 weeks of age, were immunised with identified antigen(s). The antigens include Triton X-114 extract and identified proteins of 46-48, 52-53, 60-64, 70-75, 90-94 and 110-114 kD, either singly or in combination. An immunising dose of antigen, containing between 5-100 μg protein, was given by intramuscular injection in combination with an adjuvant. An adjuvant is selected from

- (i) Seppic Montanide ISA-50
- (ii) Quill A and other derivatives of saponin,
 - (iii) oil in water emulsion employing a mineral oil such as Bayol F/Arlacel A,
 - (iv) oil in water emulsion employing a vegetable oil such as corn oil,

safflower oil or other with lecithin as emulsifier.

- (v) aluminium hydroxide gel, and
- (vi) nonionic block polymer such as Pluronic F-127 produced by BASF (U.S.A.).

Immunising doses were given at 2-4 week intervals, the number of doses being dependent on the adjuvant and amount of antigen, but preferably 2 to 3 doses are given.

Adjuvants were treated on the basis of being able to induce antibody titres, as measured by ELISA, and by assessment of induced cell-mediated immunity as tested by Delayed-Type Hypersensitivity (DTH) reaction.

The results clearly show that <u>mineral-oil type adjuvants are consistently</u> <u>superior</u> at inducing antibody titres and DTH responses (Table 2). In particular an adjuvant marketed under trade designation Montanide ISA-50 and available from Seppic, Paris, France has been found to be suitable.

TABLE 2

GROUP Anii		DTH 24 Hour	DTH 48 Hour	Antibody Levels
Nur				Lorold
	nber	Response	Response	(450 nm)
1	19	0	0	0.061
,	11	0	0	0.010
CONTROL	1	-	-	0.005
(Unvaccinated)	15	0	0	0.038
	7	0	0	0.005
•	18	+	0	0.753
QUIL A	25	+	0	0.788
1	17	0	0	0.638
1	68	- -	<u>+</u>	0.642
1	69	+++	0	0.316
2	22	0	0	0.621
VEG. OIL	4	+	0	0.666
	6	+	+	0.239
•	13	+++	++	0.457
•	14	+++	++	1.086
	5	+++	++	1.024
MIN. OIL	23	+++	+	0.864
•	16	+++	0	0.975
2	21	+	<u>+</u>	0.954

TABLE 2: Antibody levels and DTH responses in pigs measured 2 weeks after the third injection of antigen from M. hyopneumoniae. (- = no response; ± = faint reddening; + = faint reddening and swelling; ++ = reddening; +++ = swelling with or without reddening).

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Protection Pen Trial

Groups of 9 young piglets, 6 weeks of age, were immunised with purified and semi-purified antigens as shown in Table 3 below. The antigens were purified on reversed-phase HPLC using a formic acid solvent system with an acetonitrile gradient.

Antigens were resolubilised in 4 Molar urea before incorporation in mineral oil adjuvant.

The immunisation schedule is as shown in Table 2.

TABLE 3 Protocol for Pen Trial of Antigens of Mycoplasma Hyopneumoniae

5 VACCINATIONS & BLEEDS

Treatment	Day Number
1st Vaccination	0
2nd Vaccination	14
3rd Vaccination	50
Infectious Challenge	64
Slaughter	91

ANTIGEN DOSES

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Partly Purified

1st & 2nd Vaccns. 50μg COMPLEX ANTIGEN/DOSE

3rd Vaccn. - 220μg PARTIALLY PURIFIED ANTIGEN/DOSE

(Purified)74+52kD

1st Vaccn. 20μg total protein/DOSE

2nd Vaccn. 13μg total protein/DOSE

3rd Vaccn. 17μg total protein/DOSE

(Purified) 48KD 1st Vaccn. 20μg/DOSE 2nd Vaccn. 18μg/DOSE 3rd Vaccn. 27μg/DOSE

10 ALL PROTEIN ESTIMATIONS DONE BY "BCA" PROTEIN ASSAY (Pierce, Illinois, U.S.A.

Protection from infection with Mycoplasma hyopneumoniae was assessed by infectious challenge 2 weeks after the final immunisation. Infectious challenge was achieved by intranasal administration of 10ml of a 10% (w/v) lung homogenate, prepared from infected lung, and by housing test piglets with

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previously infected piglets. Four weeks after infectious challenge, the animals were killed and the extent and degree of lung lesions assessed (Table 4).

TABLE 4

Pen	Pen Trial of Antigens of Mycoplasma Hyopneumoniae					
Group No.	No. Pneumonia	Median Lung	% Reduction			
	Free (%)	Lesion Score	(from Median)			
			·			
Controls	1 (11)	13	0%			
62 kD	0 (0)	5	61%			
74+52 kD	3 (33)	6.75	48%			
48 kD	2 (22)	6.25	52%			

REFERENCE

Warren H.S. and Chedid, L.A., Future Prospects for Vaccine Adjuvants CRC Critical Reviews in Immunology 8 : 83-108, 1988.

Finally, it is to be understood that various other modifications and/or alterations may be made without departing from the spirit of the present invention as outlined herein.

CLAIMS:

1. A putative protective antigen against a <u>Mycoplasma</u>, prepared by a method including

providing

a sample of a Mycoplasma;

an antibody probe including at least one antibody against a Mycoplasma produced by a method including;

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providing a biological sample taken a short time after an immune animal has been challenged with a <u>Mycoplasma</u> or <u>Mycoplasma</u> extract taken from the infection site or an area of a lesion or an area close to the infection site or lesion;

isolating cells from the biological sample;

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culturing cells <u>in vitro</u> in a suitable culture medium; and harvesting antibodies produced from said cells;

probing the <u>Mycoplasma</u> sample with the antibody probe to detect at least one antigen; and

isolating the antigen detected.

20

- 2. A putative protective antigen according to claim 1 wherein the Mycoplasma is Mycoplasma hyopneumoniae.
- A putative protective antigen against Mycoplasma hyopneumoniae, or related infections, selected from the group of antigens having approximate molecular weights of 110-114, 90-94, 72-75, 60-64, 52-54 and 46-48 kilodaltons (kD), as herein described, mutants, derivatives and fragments thereof.
- 4. A putative protective antigen according to claim 3 which is a surface 30 protein.

- 5. A putative protective antigen according to claim 3 or 4 which is a surface lipo-protein or membrane protein.
- 6. A putative protective antigen according to any one of claims 3-5 having approximate molecular weight of 110-114, 90-94, 74, 62, 52 and 48 kD.
 - 7. A putative protective antigen according to claim 3 wherein the antigen in the 72-75 kD region contains the following N-terminal amino acid sequence:

AGXLQKNSLLEEVWYLAL

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8. A putative protective antigen according to claim 7 further including one or more of the following N-terminal amino acid sequences:

AKNFDFAPSIQGYKKIAHEL

NLKPEQILQLLG

LLKAEXNKXIEEINTXLDN

9. A putative protective antigen according to claim 3 wherein the antigen in the 60-64 kD region contains the following N-terminal amino acid sequence:

MKLAKLLKGFX(N/L)(M/V)IK

ADP(F/I)(R/E)Y(V/A)PQG(Q/A)X(M/N)VG

10. A putative protective antigen according to claim 3 wherein the antigen in the 52-54 kD region contains the following N-terminal amino acid sequence:

AGXWAKETTKEEKS

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11. A putative protective antigen according to claim 10 further including one or more of the following N-terminal amino sequences:

AWVTADGTVN

AIVTADGTVNDNKPNQWVRKY.

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12. A putative protective antigen according to claim 3 wherein the antigen in the 46-48 kD region contains the following N-terminal amino acid sequence:

AGXGQTESGSTSDSKPQAETLKHKV

13. A putative protective antigen according to claim 12 further including one or more of the following internal amino acid sequences:

TIYKPDKVLGKVAVEVLRVLIAKKNKASR AEQAITKLKLEGFDTQ KNSQNKIIDLSPEG

14. An isolated nucleic acid fragment encoding a putative protective antigen against Mycoplasma hyopneumoniae or related infections, said nucleic acid fragment including the following sequence, mutants, derivatives, recombinants and fragments thereof:

	10	20	30	40	50	
15	1234567890	1234567890	1234567890	1234567890	1234567890	
	ATGAAAAAA	TGCCACTATA	CCAGAGGAAA	GAGCAGTATA	TAAAATAATT	50
	AAAATTACAT	TTTCTTCATT	TGCGCCAGAA	TTTTTAAGAA	TTAGTACATT	100
	AAAAAGTAGA	ACAAAAGTTA	TTAATGTAAA	CATTAGCGCA	ATCCTTAAGA	150
20	AAAAATTAAA	AGTTTTATCT	ATTTTTTTA	ATCGAAATCC	AACCAGGCAT	200
	AAATCTTTGT	CAGTATTTAT	CAAGTCGGTA	TTTTTTCATT	ATTTCTACTA	250
	AAATATTATT	TGAATTTGCA	TTTTCCATAA	TCTAAAATTT	TACATTTTTT	300
	TATAACAATT	TTTAAAAATT	ACTCTTTAAT	TTATAGTATT	TTTTATTTT	350
	TTAGTCTAAA	TTATAAAATT	ATCTTGAATT	TTATTTGAAT	TTTATAATT	400
25	TAGTACTAAA	AAATACAAAT	ATTTTTCCT	ATTCTAAGAA	AAATTCATTT	450
	TTTAAAAAAA	ATTGATTTT	ATAGTATAAT	TTGTTTGTAT	AATTGAATTA	500
	ACTTGATTTG	AAAGGGAACA	AAATGAAAAA	AATGCTTAGA	AAAAAATTCT	550
	TGTATTCATC	AGCTATTTAT	GCAACTTCGC	TTGCATCAAT	TATTGCATTT	600
	GTTGCAGCAG	GTTGTGGACA	GACAGAATCA	GGTTCAACTT	CTGATTCTAA	650
30	ACCACAAGCC	GAGACGCTAA	AACATAAAGT	AAGTAATGAT	TCTATTCGAA	700
	TAGCACTAAC	CGATCCGGAT	AATCCTCGAT	GAATTAGTGC	CCAAAAAGAT	750
•	ATTATTTCTT	ATGTTGATGA	AACAGAGGCA	GCAACTTCAA	CAATTACAAA	800
-	AAACCAGGAT	GCACAAAATA	ACTGACTCAC	TCAGCAAGCT	AATTTAAGCC	850
	CAGCGCCAAA	AGGATTTATT	ATTGCCCCTG	AAAATGGAAG	TGGAGTTGGA	900
35	ACTGCTGTTA	ATACAATTGC	TGATAAAGGA	ATTCCGATTG	TTGCCTATGA	950
	TCGACTAATT	ACTGGATCTG	ATAAATATGA	TTGGTATGTT	TCTTTTGATA	1000
	ATGAAAAAGT	TGGTGAATTA	CAAGGTCTTT	CACTTGCTGC	GGGTCTATTA	1050
	GGAAAAGAAG	ATGGTGCTTT	TGATTCAATT	GATCAAATGA	ATGAATATCT	1100
	AAAATCACAT	ATGCCCCAAG	AGACAATTTC	TTTTTATACA	ATCGCGGGTT	1150
40	CCCAAGATGA	TAATAATTCC	CAATATTTTT	ATAATGGTGC	AATGAAAGTA	1200
	CTTAAAGAAT	TAATGAAAAA	TTCGCAAAAT	AAAATAATTG	ATTTATCTCC	1250
	TGAAGGCGAA	AATGCTGTTT	ATGTCCCAGG	ATGAAATTAT	GGAACTGCCG	1300
	GTCAAAGAAT	CCAATCTTTT	CTAACAATTA	ACAAAGATCC	AGCAGGTGGT	1350
	AATAAAATCA	AAGCTGTTGG	TTCAAAACCA	GCTTCTATTT	TCAAAGGATT	1400
45	TCTTGCCCCA	AATGATGGAA	TGGCCGAACA	AGCAATCACC	AAATTAAAAC	1450
	TTGAAGGGTT	TGATACCCAA	AAAATCTTTG	TAACTCGTCA	AGATTATAAT	1500
	GATAAAGCCA	AAACTTTTAT	CAAAGACGGC	GATCAAAATA	TGACAATTTA	1550

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TAAACCTGAT AAAGTTTTAG GAAAAGTTGC TGTTGAAGTT CTTCGGGTTT 1600 TAATTGCAAA GAAAAATAAA GCATCTAGAT CAGAAGTCGA AAACGAACTA 1650 AAAGCAAAAC TACCAAATAT TTCATTTAAA TATGATAATC AAACATATAA 1700 AGTACAAGGT AAAAATATTA ATACAATTTT AGTAAGTCCA GTAATTGTTA 1750 CAAAAGCTAA TGTTGATAAT CCTGATGCCT AA	TAAACCTGAT	AAAGTTTTAG	GAAAAGTTGC	TGTTGAAGTT	CTTCGGGTTT	1600
AAAGCAAAAC TACCAAATAT TTCATTTAAA TATGATAATC AAACATATAA 1700 AGTACAAGGT AAAAATATTA ATACAATTTT AGTAAGTCCA GTAATTGTTA 1750	TAATTGCAAA	GAAAAATAAA	GCATCTAGAT	CAGAAGTCGA	AAACGAACTA	1650
AGTACAAGGT AAAAATATTA ATACAATTTT AGTAAGTCCA GTAATTGTTA 1750	AAAGCAAAAC	TACCAAATAT	TTCATTTAAA	TATGATAATC	AAACATATAA	1700
	AGTACAAGGT	AAAAATATTA	ATACAATTTT	AGTAAGTCCA	GTAATTGTTA	1750
1782	CAAAAGCTAA	TGTTGATAAT	CCTGATGCCT	AA	SIMILOTIA	1782

15. An isolated nucleic acid fragment according to claim 14 encoding a putative protective antigen wherein the antigen is in the 46-48 kD region including the following nucleic acid sequence, mutants, derivatives, recombinants and fragments thereof:

	10	20	30	40	50	
	1234567890	1234567890	1234567890	. •	1234567890	
				1201007000	120-307030	
15	ATGAAAAAA	TGCCACTATA	CCAGAGGAAA	GAGCAGTATA	TAAAATAATT	50
	AAAATTACAT	TTTCTTCATT	TGCGCCAGAA		TTAGTACATT	100
	AAAAAGTAGA	ACAAAAGTTA	TTAATGTAAA	CATTAGCGCA		150
	AAAAATTAAA	AGTTTTATCT	ATTITITTA	ATCGAAATCC	AACCAGGCAT	
	AAATCTTTGT	CAGTATTTAT	CAAGTCGGTA	TTTTTTCATT	ATTTCTACTA	250
20	AAATATTATT	TGAATTTGCA	TTTTCCATAA	TCTAAAATTT	TACATTTTTT	300
	TATAACAATT	TTAAAAATT	ACTCTTTAAT	TTATAGTATT	TTTTATTTT	350
	TTAGTCTAAA	TTATAAAATT	ATCTTGAATT	TTATTTGAAT	TTTATAATT	400
	TAGTACTAAA	AAATACAAAT	ATTTTTTCCT	ATTCTAAGAA	AAATTCATTT	450
	TTTAAAAAAA	ATTGATTTTT	ATAGTATAAT	TTGTTTGTAT	AATTGAATTA	500
25	ACTTGATTTG	AAAGGGAACA	AAATGAAAAA	AATGCTTAGA	AAAAAATTCT	550
	TGTATTCATC	AGCTATTTAT	GCAACTTCGC	TTGCATCAAT	TATTGCATTT	600
	GTTGCAGCAG	GTTGTGGACA	GACAGAATCA	GGTTCAACTT	CTGATTCTAA	650
	ACCACAAGCC	GAGACGCTAA	AACATAAAGT	AAGTAATGAT	TCTATTCGAA	700
	TAGCACTAAC	CGATCCGGAT	AATCCTCGAT	GAATTAGTGC	CCAAAAAGAT	750
30	ATTATTTCTT	ATGTTGATGA	AACAGAGGCA	GCAACTTCAA	CAATTACAAA	800
	AAACCAGGAT	GCACAAAATA	ACTGACTCAC	TCAGCAAGCT	AATTTAAGCC	850
	CAGCGCCAAA	AGGATTTATT	ATTGCCCCTG	AAAATGGAAG	TGGAGTTGGA	900
	ACTGCTGTTA	ATACAATTGC	TGATAAAGGA	ATTCCGATTG	TTGCCTATGA	950
0.5	TCGACTAATT	ACTGGATCTG	ATAAATATGA	TTGGTATGTT	TCTTTTGATA	1000
35	ATGAAAAAGT	TGGTGAATTA	CAAGGTCTTT	CACTTGCTGC	GGGTCTATTA	1050
	GGAAAAGAAG	ATGGTGCTTT	TGATTCAATT	GATCAAATGA	ATGAATATCT	1100
	AAAATCACAT	ATGCCCCAAG	AGACAATTTC	TTTTTATACA	ATCGCGGGTT	1150
	CCCAAGATGA	TAATAATTCC	CAATATTTTT	ATAATGGTGC	AATGAAAGTA	1200
40	CTTAAAGAAT	TAATGAAAAA	TTCGCAAAAT	AAAATAATTG	ATTTATCTCC	1250
40	TGAAGGCGAA	AATGCTGTTT	ATGTCCCAGG	ATGAAATTAT	GGAACTGCCG	1300
	GTCAAAGAAT	CCAATCTTTT	CTAACAATTA	ACAAAGATCC	AGCAGGTGGT	1350
	AATAAAATCA	AAGCTGTTGG	TTCAAAACCA	GCTTCTATTT	TCAAAGGATT	1400
	TCTTGCCCCA	AATGATGGAA	TGGCCGAACA	AGCAATCACC	AAATTAAAAC	1450
4.5	TTGAAGGGTT	TGATACCCAA	AAAATCTTTG	TAACTCGTCA	AGATTATAAT	1500
45	GATAAAGCCA	AAACTTTTAT	CAAAGACGGC	GATCAAAATA	TGACAATTTA	1550
	TAAACCTGAT	AAAGTTTTAG	GAAAAGTTGC	TGTTGAAGTT	CTTCGGGTTT	1600
	TAATTGCAAA	GAAAAATAAA	GCATCTAGAT	CAGAAGTCGA	AAACGAACTA	1650
	AAAGCAAAAC	TACCAAATAT	TTCATTTAAA	TATGATAATC	AAACATATAA	1700
50	AGTACAAGGT	AAAAATATTA	ATACAATTTT	AGTAAGTCCA	GTAATTGTTA	1750
50	CAAAAGCTAA	TGTTGATAAT	CCTGATGCCT	AA		1782

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- 16. A method for producing an antibody against a <u>Mycoplasma</u> including providing a biological sample taken a short time after an immune animal has been challenged with a <u>Mycoplasma</u> or <u>Mycoplasma</u> extract taken from the infection site or an area of a lesion or an area close to the infection site or lesion;
 - isolating cells from the biological sample; culturing cells <u>in vitro</u> in a suitable culture medium; and harvesting antibodies produced from said cells.
- 17. A method according to claim 16 wherein the biological sample is taken at a predetermined time after the animal has been challenged with a Mycoplasma, preferably 2 to 7 days after challenge.
 - 18. A method according to claim 16 wherein the culturing of cells in vitro further includes addition of helper factors to the culture, said helper factors selected from the group including cytokines used alone or in combination, including Interleukin 1, 2, 3, 4, 5, 6, 7 and 8, colony stimulating factors, interferons and any other factors that may be shown to have an enhancing effect on specific B cell secretion.
- 20 19. A method according to any one of claims 16-18 further including a cell activation step including activating the cells isolated to proliferate and secrete and/or release antibodies

said cell activation step including adding a cell activating agent to the culture medium, said cell activating agent selected from the group including mitogens as herein described and helper factors produced by leukocytes, or their synthetic equivalents or combinations thereof.

- 20. A method according to any one of claims 16-19 wherein the antibody is in the form of the supernatant harvested from the culture medium.
- 21. An antibody against a <u>Mycoplasma</u> prepared according to the method of any one of claims 16-20.

22 .	A method	of identifying	a putative	protective	antigen	associated	with a
Myco	<u>plasma,</u> pref	ferably <u>Mycop</u>	asma hyop	<u>neumoniae</u>	, said me	ethod includi	ng
	providing						

a sample of a Mycoplasma; and

an antibody probe including at least one antibody against a Mycoplasma;

probing the <u>Mycoplasma</u> sample with the antibody probe to detect at least one antigen; and

isolating the antigen detected.

23. A method of purifying a putative protective antigen associated with a Mycoplasma, preferably Mycoplasma, said method including providing

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a crude antigen mixture; and

an antibody against a <u>Mycoplasma</u> immobilized on a suitable support;

subjecting the crude antigen mixture to affinity chromatography utilizing the immobilized antibody; and

isolating the purified antigen so formed.

- 24. A method for preparing a synthetic antigenic polypeptide against Mycoplasma, preferably Mycoplasma hyopneumoniae, which method includes providing
- 25 a cDNA library or genomic library derived from a sample of Mycoplasma; and
 - an antibody probe including an antibody prepared according to claim 16;

generating synthetic polypeptides from the cDNA library or genomic library; probing the synthetic polypeptides with the antibody probe; and isolating the synthetic antigenic polypeptide detected thereby.

- 25. A method according to claim 24 wherein the antibody probe includes an antibody raised against an antigen against Mycoplasma hyopneumoniae, or related infections, selected from the group of antigens having approximate molecular weights of 110-114, 90-94, 72-75, 60-64, 52-54 and 46-48 kilodaltons (kD), as herein described, mutants, derivatives and fragments thereof.
- 26. A synthetic putative protective antigen in the 72-75 kD region produced by a method according to claim 24 or 25 having an N-terminal amino acid sequence:

 AGXLQKNSLLEEVWYLAL

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27. A synthetic putative protective antigen according to claim 26 further including internal amino acid sequences:

AKNFDFAPSIQGYKKIAHEL
NLKPEQILQLLG
LLKAEXNKXIEEINTXLDN

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28. A synthetic putative protective antigen in the 60-64 kD region produced by a method according to claim 24 or 25 having an N-terminal amino acid sequence:

MKLAKLLKGFX(N/L)(M/V)IK

ADP(F/I)(R/E)Y(V/A)PQG(Q/A)X(M/N)VG

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29. A synthetic putative protective antigen in the 52-54 kD region produced by a method according to claim 24 or 25 having an n-terminal amino acid sequence;

AGXWAKETTKEEKS

25

30. A synthetic putative protective antigen according to claim 29 further including internal amino acid sequences:

AWVTADGTVN
AIVTADGTVNDNKPNQWVRKY.

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31. A synthetic putative protective antigen in the 46-48 kD region produced by a method according to claim 24 or 25 having an N-terminal amino acid sequence:

PCT/AU96/00149

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AGXGQTESGSTSDSKPQAETLKHKV

32. A synthetic putative protective antigen according to claim 31 further including internal amino acid sequences:

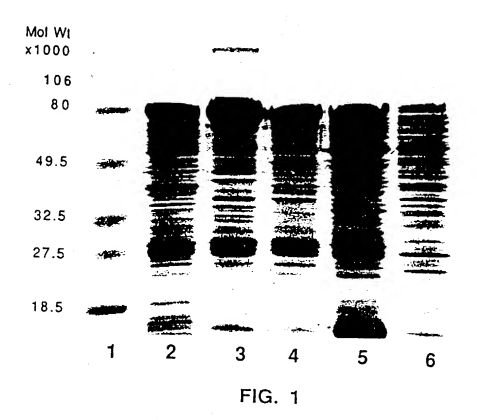
TIYKPDKVLGKVAVEVLRVLIAKKNKASR AEQAITKLKLEGFDTQ

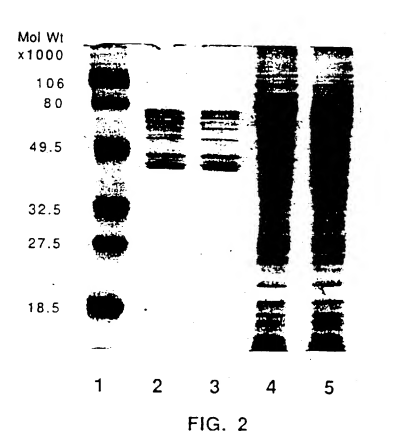
KNSQNKIIDLSPEG

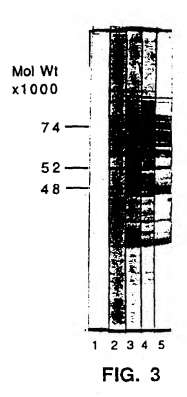
- 33. A vaccine or veterinary composition including a prophylactically effective amount of at least one putative protective antigen against a Mycoplasma according to any one of claims 1-13.
 - 34. A vaccine or veterinary composition according to claim 33 including a plurality of putative protective antigens selected from antigens having approximate molecular weights of 110-114, 90-94, 72-75, 60-64, 52-54 and 46-48 kilodaltons.
 - 35. A vaccine or veterinary composition including an antibody against a Mycoplasma according to claim 21.
 - 36. A diagnostic kit including a diagnostic antigen or fragment thereof according to any one of claims 1-13 and 26-32.
- 37. A method for preventing or treating a <u>Mycoplasma</u> infection, which method including administering to an animal a prophylactically or therapeutically effective amount of at least one putative protective antigen according to any one of claims 1-13.
- 38. An isolated DNA fragment encoding a putative protective antigen against

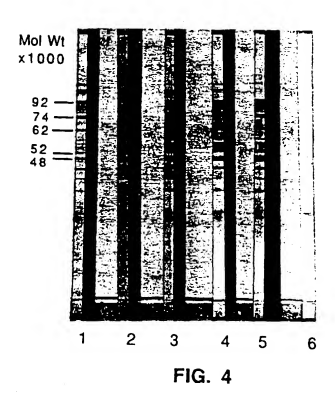
 Mycoplasma or related infections, said DNA fragment having a nucleic acid sequence according to Figure 6 or an homologous sequence, and functionally active fragments, mutant, variant or recombinant thereof.

- 39. A clone including a DNA fragment according to claim 38.
- 40. A clone according to claim 39 which is clone pC1-2 as hereinbefore described.
 - 41. An amino acid sequence or functional equivalent thereof encoded by the DNA fragment according to claim 38.
- 10 42. An amino acid sequence or functional equivalent thereof having the amino acid sequence of Figure 7.
 - 43. A putative protective antigen or antibody substantially as hereinbefore described with reference to the examples.









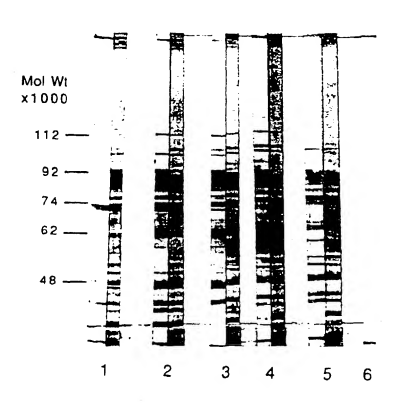


FIG. 5

			•		
10	20	30	40	50	
1234567890	1234567890	1234567890	1234567890	1234567890	
ATGAAAAAA	TGCCACTATA	CCAGAGGAAA	GAGCAGTATA	TAAAATAATT	50
AAAATTACAT	TTTCTTCATT	TGCGCCAGAA	TTTTTAAGAA	TTAGTACATT	100
AAAAAGTAGA	ACAAAAGTTA	TTAATGTAAA	CATTAGCGCA	ATCCTTAAGA	150
AAAATTAAA	AGTTTTATCT	ATTTTTTTA	ATCGAAATCC	AACCAGGCAT	200
AAATCTTTGT	CAGTATTTAT	CAAGTCGGTA	TTTTTTCATT	ATTTCTACTA	250
AAATATTATT	TGAATTTGCA	TTTTCCATAA	TCTAAAATTT	TACATTTTTT	300
TATAACAATT	TTAAAAATT	ACTCTTTAAT	TTATAGTATT	TTTTATTTT	350
TTAGTCTAAA	TTATAAAATT	ATCTTGAATT	TTATTTGAAT	TTTATAATT	400
TAGTACTAAA	AAATACAAAT	ATTTTTTCCT	ATTCTAAGAA	AAATTCATTT	450
TTTAAAAAAA	ATTGATTTTT	ATAGTATAAT	TTGTTTGTAT	AATTGAATTA	500
ACTTGATTTG	AAAGGGAACA	AAATGAAAAA	AATGCTTAGA	AAAAATTCT	550
TGTATTCATC	AGCTATTTAT	GCAACTTCGC	TTGCATCAAT	TATTGCATTT	600
GTTGCAGCAG	GTTGTGGACA	GACAGAATCA	GGTTCAACTT	CTGATTCTAA	650
ACCACAAGCC	GAGACGCTAA	AACATAAAGT	AAGTAATGAT	TCTATTCGAA	700
TAGCACTAAC	CGATCCGGAT	AATCCTCGAT	GAATTAGTGC	CCAAAAAGAT	750
ATTATTTCTT	ATGTTGATGA	AACAGAGGCA	GCAACTTCAA	CAATTACAAA	800
AAACCAGGAT	GCACAAAATA	ACTGACTCAC	TCAGCAAGCT	AATTTAAGCC	850
CAGCGCCAAA	AGGATTTATT	ATTGCCCCTG	AAAATGGAAG	TGGAGTTGGA	900
ACTGCTGTTA	ATACAATTGC	TGATAAAGGA	ATTCCGATTG	TTGCCTATGA	950
TCGACTAATT	ACTGGATCTG	ATAAATATGA	TTGGTATGTT	TCTTTTGATA	1000
ATGAAAAAGT	TGGTGAATTA	CAAGGTCTTT	CACTTGCTGC	GGGTCTATTA	1050
GGAAAAGAAG	ATGGTGCTTT	TGATTCAATT	GATCAAATGA	ATGAATATCT	1100
AAAATCACAT	ATGCCCCAAG	AGACAATTTC	TTTTTATACA	ATCGCGGGTT	1150
CCCAAGATGA	TAATAATTCC	CAATATTTTT	ATAATGGTGC	AATGAAAGTA	1200
CTTAAAGAAT	TAATGAAAAA	TTCGCAAAAT	AAAATAATTG	ATTTATCTCC	1250
TGAAGGCGAA	AATGCTGTTT	ATGTCCCAGG	ATGAAATTAT	GGAACTGCCG	1300
GTCAAAGAAT	CCAATCTTTT	CTAACAATTA	ACAAAGATCC	AGCAGGTGGT	1350
	AAGCTGTTGG	TTCAAAACCA	GCTTCTATTT	TCAAAGGATT	1400
TCTTGCCCCA	AATGATGGAA	TGGCCGAACA	AGCAATCACC	AAATTAAAAC	1450
TTGAAGGGTT	TGATACCCAA	AAAATCTTTG	TAACTCGTCA	AGATTATAAT	1500
GATAAAGCCA	AAACTTTTAT	CAAAGACGGC	GATCAAAATA	TGACAATTTA	1550
TAAACCTGAT	AAAGTTTTAG	GAAAAGTTGC	TGTTGAAGTT	CTTCGGGTTT	1600
TAATTGCAAA	GAAAAATAAA	GCATCTAGAT	CAGAAGTCGA		1650
AAAGCAAAAC	TACCAAATAT	TTCATTTAAA		AAACATATAA	1700
AGTACAAGGT	ATTATAAAAA		AGTAAGTCCA	GTAATTGTTA	1750
CAAAAGCTAA	TGTTGATAAT	CCTGATGCCT	AA		1782

1	0 20	30	40	50	
1234567890	1234567890		1234567890	1234567890	
MEEMI DEEEI	YSSAIYATSL	7 C T T 7 E777 7 C	CCOTTCCCTC	DCVDONETIV	50
	ALTDPDNPRW				100
	APKGFIIAPE	-			150
KYDWYVSFDN	EKVGELQGLS	LAAGLLGKED	GAFDSIDQMN	EYLKSHMPQE	200
TISFYTIAGS	QDDNNSQYFY	NGAMKVLKEL	MKNSQNKIID	LSPEGENAVY	250
VPGWNYGTAG	QRIQSFLTIN	KDPAGGNKIK	AVGSKPASIF	KGFLAPNDGM	300
AEQAITKLKL	EGFDTQKIFV	TRODYNDKAK	TFIKDGDQNM	TIYKPDKVLG	350
KVAVEVLRVL	IAKKNKASRS	EVENELKAKL	PNISFKYDNQ	TYKVQGKNIN	400
TILVSPVIVT	KANVDNPDA				419

FIG. 7

INTERNATIONAL SEARCH REPORT

International Application No. PCT/AU 96/00149

A. CLASSIFICATION OF SUBJECT MATTER

Int Cl6: C07K 16/12; C12N 15/31; C12P 21/00, 21/02, 21/08; G01N 33/53, 33/531; A61K 39/04; C07K 14/30

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) IPC As Above

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched AU: IPC As Above

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

WPAT, BIOT, JAPIO: MYCOPLASM: AND ANTIGEN#

CASM: MULTI-SEQUENCES

Category*	Citation of document, with indication, where	appropriate, of the relevant passages	Relevant to claim No.
х	AU,A, 70685/87 (CETUS CORP) 1 October Example 1, Claims	1987	1,16,17,20-23,33 35-37,43
x	AU,B, 49035/90 (640364) (UNIVERSITY O 11 October 1990 Claims	F MELBOURNE et al)	1,2,16,20,22
x	AU,A, 76820/91 (SYNERGEN, INC.) 17 Oc Fig. 1, 4, 6 and 7, Claim 3	tober 1991	14,38,39,41,42
x	Further documents are listed in the continuation of Box C	X See patent family annex	
"A" docum not cor "E" earlier interns "L" docum or whi anothe "O" docum exhibit "P" docum	ent defining the general state of the art which is asidered to be of particular relevance document but published on or after the ational filing date ent which may throw doubts on priority claim(s) ch is cited to establish the publication date of a citation or other special reason (as specified) ent referring to an oral disclosure, use, ion or other means ent published prior to the international filing at later than the priority date claimed	"T" later document published after the in priority date and not in conflict with understand the principle or theory us document of particular relevance; the be considered novel or cannot be considered novel or cannot be considered to particular relevance; the be considered to involve an inventive combined with one or more other sus combination being obvious to a persence." & "document member of the same pater	the application but cited to inderlying the invention e claimed invention cannot isidered to involve an taken alone e claimed invention cannot e step when the document in the documents, such on skilled in the art
Date of the actu 6 May 1996	al completion of the international search	Date of mailing of the international sear	ch r e port
AUSTRALIAN PO BOX 200	ng address of the ISA/AU INDUSTRIAL PROPERTY ORGANISATION	Authorized officer	
WODEN ACT	2606 Facsimile No.: (06) 285 3929	BARRY SPENCER	

International Application No. PCT/AU 96/00149

	tion) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
PX	AU,A, 17602/95 (SYNERGEN, INC.) 26 October 1995 Fig. 1, 4, 6 and 7 P 13 L14-P14 L5	14,38,39,41,4
x	AU,B, 49599/90 (638970) (AUSPHARM INTERNATIONAL LTD.) 26 July 1990 Examples 4 and 5	16,17,20,21,3
x	US,A, 4894332 (SCHALLER et al) 16 January 1990 Examples	1,2,14,16,17, 20-22,24,33,
x	US,A, 5252328 (FAULDS et al) 12 October 1993 Examples 1 to 4	35,37-39,41,4 1-17,20-23, 33-35,37
v	US, A, 5240706 (FAULDS) 31 August 1993	1614160
x	Examples 1 and 2, Claims EP,A, 0475185 (NIPPON FLOUR MILLS CO., LTD.)	1-6,14-16,21 22,24,25,33-3 37,38,41,42
x	Refer Example 1, Claims	1-6,12-17, 20-23,33-35 37-39,41,42
PX	Journal of Bacteriology, Vol. 177, No. 7, April 1995, pp1915-1917, "Molecular Cloning of a 46-Kilodalton Surface Antigen (P46) Gene from Mycoplasma hyopneumoniae: Direct evidence of CGG Codon Usage for Arginine", Futo et al	3-6,12-15,2 ⁴ 25,31,32,38 39,41,42
PΧ	Journal of Clinical Microbiology, Vol. 33, No. 3, March 1995, pp 680-683, "Recombinant 46-Kilodalton Surface Antigen (P46) of Mycoplasma hyopneumoniae Expressed in Escherichia Coli Can Be Used for early Specific Diagnosis of Mycoplasmal Pneumonia of Swine by Enzyme-Linked Immunosorbent assay", Futo et al	3-6, 12-15,38,39
x	Infection and Immunity. Vol. 49, No. 2, pp329-335, "surface Proteins of Mycoplasma hyopneumoniae Identified from an Esherichia coli Expression Plasmid Library", Klinkert et al	41,42 3-6,12-15,38 39,41,42
A	EP,A, 0571648 (WENG) 1 December 1993	
A	Derwent abstract Accession No. 88-010509, Class 503, JP,A, 62-273455 (NORIINSHO KK) 27 November 1987	
A	Derwent Abstract Accession No. 90-241949, Class 503, JP,A, 02-167079 (NIPPON SEIFUN KK) 27 June 1990	
A	Derwent abstract Accession No. 95-203749, Class B04, C06, D16, JP,A, 07-118167 (ZENKOKU NOGYO KYODO KUMIAI RENGOKAI) 9 May 1995	

END OF ANNEX

This Annex lists the known "A" publication level patent family members relating to the patent documents cited in the above-mentioned international search report. The Australian Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

atent Do	cument Cited in Search Report			Patent	Family Member		۸
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AU	90/49035	AT	134386	BR	9000451	CA	2008808
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		FI	900498	IL	93234	JР	3087199
	-	NO	178893	NZ	232279	ZA	9000766
AU	91/76820	AU	17602/95	CA	2078131	EP	527771
		FI	924428	HU	65827	JP	5506984
		NO	923828	wo	9115593	US	5459048
		AT	135048	AU	622855	DE	3751727
		DK	1608/88	EP	315637	HU	208550
		IL	83324	JР	1503735	wo	8800977
AU	95/17602	AU	76820/91	CA	2078131	EP	527771
		FI .	924428	HU	65827	Љ	5506984
		NO	923828	wo	9115593	US	5459048
		AT	135048	AU	622855	DE	3751727
		DK	1608/88	EP	315637	HU	208550
		IL	83324	JP	1503753	wo	8800977
AU	90/49599	EP	454735	NZ	232190	wo	9007935
		ZĄ	9000474	· · · · · · · · · · · · · · · · · · ·			
US	4894332	CA	1301677	CN	86102858	EP	196215
		JР	61274687				

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Patent Do	cument Cited in Sear Report	rch		Patent	Family Member		
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		EP	283840	HU	203672	ΙE	61626
		JР	63258427	PT	87041		
US	5240706	AT	134705	DE	68925769	EP	359919
	S	JP	2291271				
EP	475185	JP	5091882				